Exhibit 14

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92 STARCH

STARCH

Sali Bien

Starch [9005-25-8], (C₆H₁₀O₅)_n, the principal reserve polysaccharide in plants, constitutes a substantial portion of the human diet. It is the principal component of most seeds, tubers, and roots and is produced commercially from corn, wheat, rice, tapioca, potato, sago, and other sources. Most commercial starch is produced from corn which is comparatively cheap and abundant throughout the world. Wheat, tapioca, and potato starch are produced on a smaller scale and at higher prices.

Nonfood uses of starch have a long history, and Egyptian papyrus cemented with a starchy adhesive has been dated to 3500–4000 BC. Pliny the Elder (130 BC) described the uses of a modified wheat starch in sizing papyrus to produce a smooth surface (see Paper; Papermaking additives). Today, starch is used extensively in the paper, textile, and adhesive industries and has many other applications. Emergence of immobilized-enzyme technology has given rise to the production of sweeteners (qv) from corn starch (ie, corn syrup, high-fructose corn syrup), and work is underway in many laboratories to determine the extent to which cereal grains or their starches can be utilized as sources of chemical feedstocks, replacing nonrenewable petroleum sources (see Chemurgy; Enzymes, immobilized).

Starch is a mixture of linear (amylose) and branched (amylopectin) polymers of α-D-glucopyranosyl units. Natural starch occurs usually as granules composed of both linear and branched starch molecules. However, some starches are composed only of branched molecules, and these are termed waxy starches because of the vitreous sheen of a cut seed surface. Some mutant seed varieties have been produced with starches having up to 85% linear molecules, although most starches have ca 25% linear and 75% branched molecules.

The quasi-crystalline granules that characterize starch ultrastructure are insoluble in water at ambient temperature and relatively resistant to carbohydrases other than α-amylases (see also Carbohydrates; Sugar).

Physical Properties

Starch occurs in plants in the form of granules which may vary in diameter from 2-150 µm (1). Rice starch has the smallest granules and potato starch the largest (2). Microscopic examination of starch granules reveals a distinct cleft called the hilum granule growth occurred. Microscopic examination with polarized light reveals a birefringence which, along with x-ray diffraction, is evidence of semicrystallinity of the granule. Between crossed Nicol prisms of the microscope, a black cross, ie, a cross of isoclines, is observed at the center of the hilum. Cereal starches give an A-type x-ray diffraction, tuber starches a B-type pattern, and a few starches give an intermediate diffraction pattern, the C type (3).

Undamaged starch granules are insoluble in cold water but imbibe water reversibly and swell slightly. The percent increase in granule diameter ranges from 9.1% for corn to 22.7% for waxy corn (4). However, in hot water, a larger irreversible swelling occurs producing gelatinization. Gelatinization takes place over a discrete temperature range that depends on starch type.

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 Starch
 Range, °C

 potato
 59-68

 tapioca
 58.5-70

 corn
 62-72

 waxy corn
 63-72

 wheat
 58-64

peting for available water. Other chemicals, such as sodium nitrate, alkali, and urea, inhibit gelatinization and increase the gelatinization temperature, probably by comapplications of starch. Certain chemicals, such as sodium sulfate, sucrose, and dextrose, range in a predictable way, and this information may be important to certain industrial a Kofler hot stage. Other chemicals present in the slurry may affect the gelatinization molecular hydrogen bonds. lower the gelatinization temperature range, possibly by disrupting granular intermethod follows microscopically the loss of birefringence of a starch slurry heated on gelatinization range depends upon the method used to measure it. The most sensitive limit, corresponding to the point where almost all granules are 100% gelatinized. The is the range between the lower limit, indicating onset of granule swelling, and the upper thick pastes or gels is the basis of many starch uses. The gelatinization temperature thickens and, if sufficiently concentrated, may form a gel. The property of forming amylose molecules, swollen granules, and granule fragments is cooled, the mixture are leached out of the granule into solution. When a cooked starch paste containing size. During swelling, and as a consequence of it, some of the linear amylose molecules are solvated first and the granule swells rapidly, eventually to many times its original bonding in the interior of the starch granule. The amorphous regions of the granule temperature), the kinetic energy of the system is sufficient to overcome the hydrogen At a certain temperature in the heating process (the lower limit of the gelatinization

The physical properties of starch are altered by mechanical treatment. If in the dry state the granular integrity is disrupted, as by grinding, the starch gelatinizes more readily in cold water. Furthermore, the granule is readily attacked by chemicals and enzymes. In the swollen state, the granule is more fragile, and vigorous agitation of a cooked paste results in rupture of most, if not all, granules. Consequently, the cold paste loses its viscosity and gelling ability.

Specific optical rotation values $[\alpha]_D$ for starches range from 180 to 220° (3); for pure amylose and amylopectin fractions, $[\alpha]_D$ is 200°. Measurements of these values are hampered by the limited solubility of these polymers and the opacity of some of their dispersions. More recently, the structure of amylose has been investigated with infrared spectroscopy (6). Analysis of the spectrum shows it to be consistent with the proposed ground-state conformation of the monomer D-glucopyranosyl units. The nature of intramolecular bonding in amylose has also been investigated with nmr spectroscopy (7).

Chemical Properties

Most common starches contain two different types of D-glucopyranose polymers. Amylose is essentially a linear polymer of α -D-glucopyranosyl units linked (1--4) as shown in Figure 1. This polymer may be separated from the starch by complete gelatinization and vigorous dispersion of the hot starch solution with a complexing agent such as 1-butanol (8) in water. On cooling, an amylose-butanol complex crystallizes

fabric finishes (see Textiles). In this application they serve to increase yarn strength and abrasion resistance and generally improve weaving efficiency. Thin-boiling starches also find some applications in paper manufacture and laundry-starch prep-

Oxidized starches, principally those prepared by hypochlorite oxidation, are mainly used in paper coatings and adhesives to improve surface characteristics for mainly used in paper coatings and adhesives to improve as textile warp sizes and printing or writing. Oxidized starches are also employed as textile warp sizes and finishes, in the manufacture of insulation and wallboard, and in laundry spray

Starch pyrodextrins and British gums have the ability, in aqueous solution, to Starch pyrodextrins and British gums have the ability, in aqueous solution, to form films capable of bonding like or unlike materials. As such, they have extensive use as adhesives for envelopes, postage stamps, and other products (see Adhesives). These dextrins are used in glass-fiber sizing, to protect the extruded fiber from abrasion, and as binders for metal core castings, water colors, briquettes, and other com-

posite materials.

Various organic chemicals, eg, ethanol, isopropyl alcohol, n-butanol, acetone, Various organic chemicals, eg, ethanol, isopropyl alcohol, n-butanol, acetone, 2,3-butylene glycol, glycerol, and fumaric acid, are produced from starch by fermen-2,3-butylene glycol, glycerol, other polyhydroxy compounds from starch include tation (74) (see Fermentation). Other polyhydroxy compounds from starch include tation (74) (see Fermentation). Other polyhydroxy compounds from starch include tation (74) (see Fermentation).

pyranosiues.

Food Uses. Unmodified starch is used in food preparations requiring thickening, food Uses. Unmodified starch is used in food preparations shall dressings, gelling, or similar properties. Such applications include puddings, salad dressings, gelling, and candies. Pre-gelatinized starch is used in a variety of products where pie fillings, and candies. Pre-gelatinized starch is used in a variety of products where thickening is required but cooking is to be avoided, such as instant pudding, pie fillings,

and cake frostings.

Acid-modified starches are used for the manufacture of gum candies because they Acid-modified starches are used form hot concentrated pastes that gel firmly on cooling. Heat-treated starches are used in food applications to bind and carry flavors and colors. Sweetening agents are made from starch by enzymatic or acid treatment.

Derivatives

Starches, as polyhydroxy compounds, undergo many reactions characteristic of alcohols, including esterification and etherification. Since the D-glucopyranosyl monomers contain, on average, three free hydroxyl groups, the degree of substitution (DS) may be at most three. The more important commercial starch derivatives are very lightly derivatized (DS <0.1). Such modifications may produce distinct changes in colloidal properties and generally produce polymers with properties useful in a wide

variety of applications.

Hydroxylalkyl Starch Ethers. Hydroxyethyl starch ethers of low DS (0.05-0.10) have been produced in a number of ways, usually near the end of the wet-milling process on a high solids-starch suspension. The derivative, a modification of ungelatinized starch, is easily filtered and can be produced economically in a fairly pure form

(see also Cellulose derivatives).
In the wet-milling process, a 40–50% solids-starch suspension is treated with a In the wet-milling process, a 40–50% solids-starch suspension is treated with a Group I or Group II metal hydroxide and ethylene oxide at a 50°C. A DS of 0.1 is easily obtained, and the product is still conveniently purified by filtration. Higher DS products are obtainable but are more difficult to purify and filter because the derivatized granules swell during washing.

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Introduction of hydroxyethyl groups at low DS results in marked modification of physical properties. Among them are reduced gelatinization temperature (75), increased rate of granule swelling (76), and lowered tendency of starch pastes to gel and retrograde.

Low DS hydroxyethyl starches are widely used as paper coatings and sizes to improve sheet strength and stiffness. They are also used as paper coating color adhesives and to increase internal fiber bonding in paper products. Additionally, hydroxyethyl starches are used as textile warp sizes.

Hydroxypropyl and other hydroxylalkyl starches are used as additives in salad dressing, pie fillings, and other food thickening applications (see Food additives).

dressing, pie fillings, and other took structures are the tertiary and Cationic Starches. The two commercial cationic starches are the tertiary and Cationic Starches. The two commercials cationic starches are prepared by treating quaternary aminoalkyl ethers. Tertiary aminoalkyl ethers are prepared by treating a basic starch slurry with a tertiary amine containing a β-halogenated alkyl, 3- a basic starch slurry with a tertiary amine containing a β-halogenated alkyl, 3- a basic starch slurry with a tertiary amine free bases. Treatment starch ethers are readily formed that contain tertiary amine free bases. Treatment with acid gives the cationic form. Amines that have been used in this reaction include with acid gives the cationic form. Amines that have been used in this reaction include 2-dimethylaminoethyl chloride, 2-diethylaminoethyl chloride, and M-(2,3-epoxy-popy)] diethylamine. Commercial preparation of low DS derivatives requires reaction propally the start of the start o

Cationic starches exhibit decreased gelatinization temperature and increased hot paste viscosity. Pastes remain clear and fluid even at room temperatures and show no tendency to retrograde. This stability is undoubtedly due to repulsion between the cationic starch molecules in dispersion.

Quaternary ammonium alkyl ethers are prepared in similar fashion: an alkaline starch reacts with a quaternary ammonium salt containing a 3-chloro-2-hydroxypropyl or 2,3-epoxypropyl radical. Alternatively, these derivatives may be prepared by simple quaternization of tertiary aminoalkyl ethers by reaction with reagents such as methyl iodide. Sulfonium (77) and phosphonium (78) starch salts have also been prepared

and examined.

Quaternary ammonium starches, like the tertiary ammonium derivatives, show Quaternary ammonium starches, increased paste clarity and viscosity, and redecreased gelatinization temperatures, increased paste clarity and viscosity, and redecreased gelatinization temperatures, increased paste clarity and viscosity, and redecreased gelatinization temperatures are used on paper mainly for fiber swelling at a DS as low as 0.07. Cationic starches are used on paper mainly for fiber swelling at a DS as low as 0.07. Cationic starches are used in the paper sizes. endurance. They have been employed as emulsifiers for water-repellent paper sizes, although they have been employed in ore refining as flocculating agents (qv).

Starch Phosphates. Starch phosphate monoesters may be prepared by heating a dry mixture of starch and acid salts of ortho-, pyro-, or tripolyphosphoric acid at a dry mixture of starch and acid salts of ortho-, pyro-, or tripolyphosphoric acid at 50-60°C for 1 h. Degree of substitution is generally low (DS <0.15), but higher DS derivatives can be prepared by increasing the temperature, phosphate salt concentration, and reaction time.

Compared to the unmodified starches, the monophosphate esters have a lower Compared to the unmodified starches, the monophosphate esters have a lower gelatinization temperature and swell in cold water at a DS of 0.07. Like other derivatives, starch phosphates have increased paste viscosity and clarity and decreased set-back or retrogradation. Their properties are similar to those of potato starch, which also contains phosphate groups.

in aluminum refining from bauxite ores, and as detergent builder. in papermaking to improve fold strength and surface characteristics, as a textile size, powders and icings. Nonfood applications include use as core binders for metal molds, has been developed (79) which is dispersable in cold water for use in instant dessert preparations, they are superior to other starches. A pregelatinized starch phosphate excellent freeze-thaw stability. As thickeners in frozen gravy and frozen cream-pic Starch monophosphates are very useful in food applications because of their

is more resistant to heat, agitation, acid, and rupture of swollen granules. granule produces a starch which, compared to the unmodified starch, swells less and between two or more starch chains. The covalent linkage of the starch chains in the In contrast to monophosphates, starch phosphate diesters contain ester cross-links

a starch slurry and 2% trimetaphosphate salt react at pH 10-11 and 50°C for 1 h. (81), which require more vigorous conditions than phosphorus oxychloride. Typically, phorus oxychloride, phosphorus pentachloride, and thiophosphoryl chloride (80) Cross-bonded starches may also be produced by the reaction with trimetaphosphates Starch in aqueous suspension may be cross-linked to form diesters with phos-

in boiling water. greater or less than that of the parent starch. In contrast to starch phosphate mogranules. Depending on the degree of derivatization, the hot paste viscosity may be If the degree of cross-linking is sufficient, swelling can be completely inhibited, even noesters, the pastes of the diesters do not increase in clarity. Starches with high DS nave exceptional stability to high temperatures, low pH, and mechanical agitation. Starch phosphate diesters show a significant increase in the stability of the swollen

on standing. They are also used to produce high wet-strength paper (80), as ion-exresistance to gelling and retrogradation, freeze-thaw stability, and lack of syneresis unmodified starches in their ability to keep food in suspension after cooking, increased foods, salad dressings, fruit pie filling, and cream-style corn. They are superior changers, and metal sequesterants to prevent oxidative rancidity in oils. Cross-linked starch derivatives are used as thickeners and stabilizers in baby

sols. Light derivatization of low DS acetate inhibits association of the amylose polymers solubility in organic solvents and ability to form films and fibers. applications. More highly derivatized starches (DS 2–3) are useful because of their and the longer outer chains of amylopectin. These properties are important in food importance of low DS acetates is based mainly on the stabilization of aqueous polymer Starch Acetates. Starch acetates may be either low DS or high DS. The commercial

dride to make low DS starch acetates; ketene or vinyl acetate have also been investiacid or preferably acetic anhydride, either alone or in acetic acid, pyridine, or aqueous 7-11 and room temperature to give a DS of 0.5. gated. Commercially, the acetic anhydride-aqueous alkali system is employed at pH alkaline solution. Dimethyl sulfoxide has been used as a cosolvent with acetic anhy. Low DS starch acetates are produced by treatment of granular starch with acetic

paper manufacture. is high enough. Low DS starch acetate polymers also form films useful in textile and to retrograde after pasting and cooling. Gelling may be completely inhibited if the DS Low DS starch acetates have lower gelatinization temperatures and less tendency

in paper to improve printability, surface strength, and solvent resistance. stability. These applications include frozen fruit pies and gravies, baked goods, instant puddings, and pie fillings. Starch acetates are employed in textiles as warp sizes and lightly derivatized starch acetates are used in food because of their clarity and

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acetates give weak and brittle films and fibers. However, amylose triacetate is useful but with longer reaction time. In general, high DS starch acetates and amylopectin pliable, lustrous, transparent, and colorless. These properties are useful in packaging and chloroform. Films of this high DS acetate, cast from chloroform solution, are in forming film and fiber. It is soluble in organic solvents such as acetic acid, pyridine High DS starch acetates are prepared by the methods used for low DS acetates

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